



## UNITED STATES DEPARTMENT OF COMMERCE Patent and Trademark Office

Address: COMMISSIONER OF PATENTS AND TRADEMARKS Washington, D.C. 20231

SERIAL NUMBER FILING DATE FIRST NAMED APPLICANT ATTORNEY DOCKET NO. 04724787 07/042,498 **ULBELLE** SCULLY, SCOTT, MURPHY & PRESSER HOPFERSE 200 GARDEN CITY PLAZA GARDEN CITY, NY 11530-3391 ART UNIT PAPER NUMBER DATE MAILED: 04/12/88

This is a communication from the examiner in charge of your application. COMMISSIONER OF PATENTS AND TRADEMARKS

This	s application has been examined	Responsive to communication file	1 on	This action is made final.	
A shortened statutory period for response to this action is set to expire month(s), days from the date of this letter.  Failure to respond within the period for response will cause the application to become abandoned. 35 U.S.C. 133					
Part I  L  3. [ 5. [	THE FOLLOWING ATTACHMENT Notice of References Cited by Ex Notice of Art Cited by Applicant, Information on How to Effect Draw	PTO-1449 <b>4.</b>	Notice re Patent Drawing Notice of informal Paten	g, PTO-948. t Application, Form PTO-152	
Part II SUMMARY OF ACTION					
ار ۱۰	Claims 1-10		•	are pending in the application.	
	Of the above, claims			are withdrawn from consideration.	
2. [	Claims			_ have been cancelled.	
3. [	Claims			_ are allowed.	
4. j	Claims 1-10			are rejected.	
5. [	Claims	4-7-11-11-11-11-11-11-11-11-11-11-11-11-1		_ are objected to.	
6. [	Claims	Claims are subject to restriction or election requirement.			
7.		This application has been filed with informal drawings which are acceptable for examination purposes until such time as allowable subject matter is indicated.			
8. [		Allowable subject matter having been indicated, formal drawings are required in response to this Office action.			
9. [		The corrected or substitute drawings have been received on These drawings are acceptable; not acceptable (see explanation).			
10. [		The proposed drawing correction and/or the proposed additional or substitute sheet(s) of drawings, filed on has (have) been approved by the examiner disapproved by the examiner (see explanation).			
11. [	the Patent and Trademark Office corrected. Corrections MUST be	The proposed drawing correction, filed, has been approved disapproved (see explanation). However, the Patent and Trademark Office no longer makes drawing changes. It is now applicant's responsibility to ensure that the drawings are corrected. Corrections MUST be effected in accordance with the instructions set forth on the attached letter "INFORMATION ON HOW TO EFFECT DRAWING CHANGES", PTO-1474.			
12. [	Acknowledgment is made of the c	Acknowledgment is made of the claim for priority under 35 U.S.C. 119. The certified copy has been received not been received			
		been filed in parent application, serial no; filed on			
13. [	Since this application appears to be in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under Ex parte Quayle, 1935 C.D. 11; 453 O.G. 213.				
14. [	Other				

- 1. The Abstract of the Disclosure is objected to because if fails to fully comply with the requirements under 37 CFR 1.72(b). The abstract should be included under the heading --Abstract of the Disclosure-- and not merely "Abstract". Correction is required. See MPEP 608.01(b).
- The disclosure is objected to because of the following informalities:

In the specification on page 20, line 6, it is uncertain if the author "Mulskin" is intended or if --Milstein-- is intended.

Appropriate correction of the disclosure is required.

- 3. Claims 1-4 are rejected under 35 U.S.C. 112,
  first paragraph, as the disclosure is enabling only for
  claims limited in accordance with the specification.
  The broad term "antigen" is considered to be enabled by
  an antigen of Ovarian cancer as set forth in the speciis only enabled for the specific type of antigen recited therein and not enabled for any
  fication. The antigen/with a 40 kilodalton molecular
  weight, therefore, the claims lack enablement in the
  specification. See MPEP 706.03(n) and 706.03(z).
- 4. Claims 1-10 are rejected under 35 U.S.C. 112, second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which applicant regards as the invention.

The recital of "CA125" renders claim 1 ( and all other instances) vague and indefinite as to the intended meaning. Is the CA125 antigen present only in CVACIAN cancer? The recitation of "time sufficient" renders claim 6 vague and indefinite as to the intended meaning. What do Applicants consider a sufficient time for the formation of a binary complex?

5. 35 U.S.C. 101 reads as follows:

Whoever invents or discovers any new and useful process, machine, manufacture, or composition of matter or any new and useful improvement thereof, may obtain a patent therefor, subject to the conditions and requirements of this title.

- 6. Claims 1 and 2 are rejected under 35 U.S.C.

  101 because the claimed invention is directed to nonstatutory subject matter. Since the claims do not specifically recite "an isolated substantially purified
  subunit of CA125 antigen" they read on an antigen that
  exists in its native state, in vivo and are thus nonstatutory.
- 7. The following is a quotation of the appropriate paragraphs of 35 U.S.C. 102 that form the basis for the rejections under this section made in this Office action:

A person shall be entitled to a patent unless-

8. (b) the invention was patented or described in a printed publication in this or a foreign country or in public use or on sale in this country, more than one year prior to the date of application for patent in the United States.

9. The following is a quotation of 35 U.S.C. 103 which forms the basis for all obviousness rejections set forth in this Office action:

A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.

Subject matter developed by another person, which qualifies as prior art only under subsection (f) and (g) of section 102 of this title, shall not preclude patentability under this section where the subject matter and the claimed invention were, at the time the invention was made, owned by the same person or subject to an obligation of assignment to the same person.

10. Claims 1-4 are rejected under 35 U.S.C. 102
(b) as anticipated by or, in the alternative, under 35
U.S.C. 103 as obvious over Masuno et al and Bast et al.

Masuno et al discloses a monoclonal antibody,

to

OC 125, specific Adisinct determinants on the surface of
human CNACIAN cancer cells. Bast et al disclose

monoclonal antibody, OC125, which recognizes the antigen

CA125 which is the same antigen of the present invention. It is noted that Bast et al does not disclose a

particular submit of this antigen. The specific determinants and antibody of Masuno et al inherently have the
same characteristics as required by the claims. Since

the antigens are identical they would inherently posses the same specific cent-surface determinants or submits. A difference in molecular weight could result from a difference in purity and/or a difference in molecular weight determination. In the event of unexpected results it would have been well within the purview of the skilled artisan through limited routine experimentation to detect the claimed subunit with highly conventional techniques

11. Claims 5-10 are rejected under 35 U.S.C. 103
as being unpatentable over Masuno et al and Bast et al inview of Pestka or the WO Patent.

Masuno et al and Bast et al disclose as set
forth supra. Pestka and the WO Patent disclose a twosite or sandwich assay employing polyclonal and
mononclonal antibodies. A two-site assay is used to
detect two distinct epitopesor determinants at two different sites on an antigen.

When detecting for the specific determinants of the CAl25 antigen with the monoclonal antibodies of Masuno et al and Bast et al it would have been obvious to one of ordinary skill in the art to use the two-site or sandwich assay of Pestka or the WO Patent for the advantages associated with such assay methods. Also, the formation of a kit would have obvious since such kits are highly conventional.

The prior art made of record and not relied upon is considered pertinent to applicant's disclosure.

Mattes et al disclose diagnosis and therapy for cancer patients employing monoclonal antibodies to several cell antigens of human OVACIAN, cervical and endemetrial carcinomas.

Knauf discloses a monoclonal antibody to a specific antigen derived from human ovarian carcinomas.

Also disclosed is the use of this antibody in a radioimmuno assay for the detection of evarian carcinomas.

The prior art reference cited by applicant on page 4 of the specification (Knaaf and Urback, AM. J. Obstet. Gynecol. 138: 1222, 1980) is not readily available to the Examiner. Applicants assistance is providing a copy to complete the record would be appreciated.

Any inquiry concerning this communication should be directed to Florina B. Hoffer at telephone number 703-557-0664.

HOFFER/fm

4/3/88

ROBERT J. WARDEN
SUPERVISORY PATENT EXAMINER
ART UNIT 128

Robert Y. Wandon